



















































Table 1: Epilepsy Syndromes and conditions (listed	the KD has been reported moderately beneficial (not better
alphabetically) for which the KD has been consistently	than the average dietary response, or in limited single-
reported as more beneficial (>70%) than the average 50% KD	center case reports)
response (defined as >50% seizure reduction).	Adenylosuccinate lyase deficiency <sup>64</sup>
Angelman syndrome <sup>56,57</sup>	CDKL5 encephalopathydisorders <sup>67</sup>
Complex 1 mitochondrial disorders <sup>51,55</sup>	Childhood absence epilepsy <sup>69</sup>
Dravet syndrome <sup>35,36</sup>	Cortical malformation <sup>73, 74</sup>
Epilepsy with myoclonic-atonic seizures (Doose	Epilepsy of infancy with migrating focal seizures <sup>68</sup>
syndrome) <sup>34,37,38</sup>	Epileptic encephalopathy with continuous spike-and-wave
Glucose transporter protein 1 (Glut-1) deficiency syndrome	during sleep <sup>70</sup>
(Glut1DS) <sup>27,29-32</sup>	Glycogenosis type V <sup>65</sup>
Febrile infection-related epilepsy syndrome (FIRES) <sup>44-47</sup>	Juvenile myoclonic epilepsy <sup>bb</sup>
Formula-fed (solely) children <sup>48,49</sup>	Latora body disease <sup>36</sup>
Infantile spasms <sup>10,39,40</sup>	
Ohtahara syndrome <sup>50–52</sup>	Lennox-Gastaut syndrome <sup>26</sup>
Pyruvate dehydrogenase deficiency (PDHD) <sup>28</sup>	Phosphotructokinase deficiency <sup>03</sup>
Super-refractory status epilepticus <sup>44,46,53,54</sup>	Rett syndromess, or
luberous scierosis complex <sup>41–43</sup>	Subacule scierosing pariencephantis (SPSE)
*full references available in article	*full references available in article
Patients who fail the traditional anti-con	vulsant therapy
poor candidates for epilepsy surgery	
All ages	
Average time on diet 1-2 years	
12020 Nutricia North America	Tables adapted from Kossoff et al. Epilensia Open. 2018;3:175.
2020 Nutricia North America	lables adapted from Kossoff, et al. Epilepsia Open. 2018;3:175



	Table 3: Contraindications to the use of the KD			NK
	Absolute			UN
	Carnitine deficiency (primary)			
	Carnitine palmitoyl transferase (CPT) I or II deficiency			
	Carnitine translocase deficiency			
	β-oxidation defects			
	Medium-chain acyl dehydrogenase deficiency (MCAD)			
	Long-chain acyl dehydrogenase deficiency (LCAD)			
	Short-chain acyl dehydrogenase deficiency (SCAD)			
	Medium-chain 3-hydroxyacyl-CoA deficiency			
	Pyruvate carboxylase deficiency			
	Porphyria			
	Relative			
	Inability to maintain adequate nutrition			
	Surgical focus identified by neuroimaging and video-EEG monitoring			
	Parent or caregiver noncompliance			
	Propofol concurrent use (risk of propofol infusion syndrome may be h	nigher)		
utricia N	lorth America	Table adapted from Kossoff,	et al. Epilepsia Open. 2018;3:17!	5-92.























Ratio	Calories per Dietary Unit	Example
2:1	22	(2X9) + (1X4) =22
3:1	31	(3X9 )+ (1X4) =31
4:1	40	(4X9 )+ (1X4) =40
5:1	49	(5X9) +(1X4)=49
By dividing the calories (age x and fat in a giv	e dietary units of a given ratio calories per kg) the total gran en ketogenic diet can be dete	into the determined t ns of protein, carbohy ermined.



Day	Topics Covered	
	MD review	
	Diet basics	
Monday	Side Effects	
	Diet initiation schedule	
	Meet with floor team/nurses	
	RD: The basics of the ketogenic diet	
Tuesday	Meal Plan Guidelines	
	Social worker meeting	
	Ketogenic Computer program	
	<ul> <li>What to do when your child gets sick</li> </ul>	
Wednesday	Parent lecture	
weathesday	Weighing and measuring foods	
	Learning how to read recipes	

# Side Effects

- Constipation
- Poor growth
- Osteopenia/ osteoporosis
- Kidney stones
- Hyperlipidemia

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• Vitamin and mineral deficiency

Follow-up	Table 6: Recommendations for aspects of a follow-up KD clinic visit           Nutritional assessment (registered dietitian)
<ul> <li>Generally every 3 months for the initial year</li> <li>Children less than 2 years seen in 1 month</li> <li>Labs at each visit <ul> <li>CMP, Fasting lipid panel, urinalysis, Vit D, selenium</li> </ul> </li> </ul>	Height, weight, ideal weight for stature, growth velocity, BMI when appropriate         Head circumference (when appropriate)         Review appropriateness of KD prescription (calories, protein, and fluid)         Review vitamin and mineral supplementation       Assess compliance to KD         Adjust KD, if necessary, to improve compliance and seizure control         Medical evaluation (neurologist)         Efficacy of the diet (is the KD meeting parental expectations?)         Side effects of the KD       Antiseizure drug reduction
Medications usually not changed the first month	applicable) Should KD be continued? Laboratory assessment Complete blood count with platelets
Frequent phone and email contact in between clinic visits	Electrolytes to include serum bicarbonate, total protein, calcium Serum liver and kidney profile (including albumin, blood urea nitrogen, creatinine) Vitamin D level Free and total carnitine Selenium level SEG(Critical Selenium level SEG(Critical Selenium level
At our center, all management through the keto team from now on	Urinaysis EEG (at KD discontinuation) Anticonvulsant drug levels (if applicable) Optional Serum beta-hydroxybutyrate (BOH) level Urine calcium and creati
Table adapted from Kossoff, et al. Epilepsia Open. 2018;3:175-92.	Zinc, copper levels Renal ultrasound ECG (Electrocardiogram) Bone mineral density (DEXA scan) after 2 years on the KD *Visits should be at least every 3 months for the first year of the KD, with a visit 1 mo after starting the KD also advised

# Other Diet Variants for Epilepsy Modified Atkins Diet Low Glycemic Index Diet MCT oil Diet Modified Ketogenic Diet

Calories not restricted, based on RDA's or individual needs

39

### Comparison KETO Modified Atkins Diet Low Glycemic Index Diet No calorie or protein restriction Focus on low glycemic index carbohydrates <50 No weighing on gram scale Allowed a lot more carbohydrates Stress the importance of fat in the diet **Modified Ketogenic Diet** MCT Oil Diet Uses medium chain triglycerides (MCT oil) in place of the large • Low and slow amounts of cream & butter to allow more room for carbs and Start off at a low ratio and slowly increase as needed • protein Based on percentages of calories vs. ratio's (ex. 30-60% MCT oil, 10% carb, 12% protein or 2 x RDA, 18-48% fat). Total fat=70-80%\*

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**KETO** 

# Johns Hopkins Adult Epilepsy Diet Center: Modified Atkins Diet Protocol

### **Pre-Diet**

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- Nutrition evaluation 3-day food record, anthropometrics
- Laboratory values CMP, CBC, fasting lipids
- Medications levels
- Diagnostic studies: EEG, MRI..
- Screening for cardio and cerebrovascular risk factors, history of kidney stones

## Initiation

- 20 g of carbohydrates
- No calorie restriction
- MVI, Calcium and Vitamin D
- Seizure calendar
- Urine ketones
- Weight
- Start and end of menses







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